Attachment 1

Piscivory by Non-Native Salmonids in the Colorado River and an Evaluation of the Efficacy of Mechanical Removal of Non-Native Salmonids

An Operational Plan

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EXECUTIVE SUMMARY

Need

The recommended flows contained within the treatment scenarios related to testing fish hypotheses center around the notion of improving future humpback chub (HBC) recruitment by reducing the number of adult rainbow trout (RBT) and brown trout (BNT) residing in the system downstream of Lee's Ferry. This study will address questions raised by the Technical Working Group (TWG) of the Glen Canyon Adaptive Management Program asking 1) whether or not reducing RBT and BNT abundance will improve HBC recruitment and a related question 2) are RBT and BNT significant predators of HBC? This study will also address a number of issues identified by the aquatic protocol evaluation panel (Anders et al. 2001). The panel had concerns with the lack of empirically established linkages between food base and fishes, and identified that a possible consequence of the recent increase in primary and secondary production may differentially benefit non-native species (competitors or predators) over native species. Secondly, the panel identified the need for establishing a better understanding of the relationship and trophic linkages between foodbase and fish. Therefore, the trout dietary analysis in this study will be integrated with other existing GCMRC long-term monitoring programs that are presently collecting or proposing to collect data specific to: 1) aquatic benthic foodbase, and 2) carbon productivity monitoring program.

Benefits

With the potential removal of non-native fishes that may prey on or compete with humpback chub, recruitment of humpback chub may increase. This study will determine if mechanical removal of salmonids is feasible in a large river ecosystem. The mechanism for possible increased humpback chub recruitment will be determined through diet analyses of salmonids. In addition, diet analyses will determine the size range of piscivorous salmonids and the relationship between prey size and predator size.

Objectives

The objectives of this study are to determine 1) the efficacy of mechanical removal of adult RBT and BNT from the LCR Inflow reach, 2) RBT and BNT predation and diet, and 3) the effect of adult RBT and BNT in the LCR inflow reach on the population dynamics of the LCR HBC population. In addition, we will continue to monitor the HBC population downstream of the LCR to determine mortality/emigration rates from the LCR reach.

Study Area

We have selected a study area in the Colorado River (56.2 RM - 65.7 RM) that encloses the majority of the geographic distribution of the Little Colorado River humpback chub population. The upstream and downstream study area endpoints are bounded by hydraulic and geomorphic controls (Kwagunt and Lava Chuar Rapid). A control reach has also been established between RM 44 and RM 52 (President Harding Rapid to Nankoweap).

Procedures

We will conduct annually, three depletion trips in January-March and three depletion trips in July-September. The annual depletion efforts will be repeated four years, for a total of 24 times, to determine how removal of fish using a series of depletion passes in a discrete area will influence the relative abundance of the remaining fish stock. The sampling efforts are scheduled to coincide with the major periods of LCR flooding events (spring runoff and monsoonal storms) that are correlated with juvenile HBC immigration to the mainstem Colorado River (Valdez and Ryel 1995). Non-native fishes will be collected, euthanized, and disposed. All native fishes will be measured, weighed, tagged, and released. Stomach contents will be collected from all non-native fish to determine incidence of predation on humpback chubs. To determine prey selection for invertebrates by salmonids, entire stomach contents from a sub-sample of non-native fish will be collected and compared with invertebrate drift samples collected during the same time as the fish collections.

Deliverables

Semi-annual reports and presentations will be given to the Adaptive Management Work Group and /or the Technical Work Group during December and June of each year of the study. At least four peer-reviewed publications in the

primary literature are expected from this body of work. The anticipated submittal date for these peer-reviewed publications is 2004-2006.

INTRODUCTION

Background

Recent analyses of historical humpback chub (HBC) data suggest that the abundance of the Little Colorado River (LCR) population is in decline (Figure 1; Grand Canyon Monitoring and Research Center (GCMRC) unpublished analyses). These analyses utilized mark-recapture data in an open population model to construct estimates of the population recruitment (1989-1998 brood years) and sub-adult and adult abundance (>150 mm total length; 1991-2000). The decline in the abundance of sub-adult and adult fish appears to be the result of continued low recruitments beginning with the 1992 brood year. As these weak year classes have entered the sub-adult and adult portions of the population, the overall abundance of HBC has declined from a peak of 8,279 in 1993 to 2,515 in 2000. The overall trends in recruitment and abundance are supported by two additional analyses. First, the downward recruitment trend is supported by trends observed in the catch-rate (CPUE) of Age-1 and Age-2 HBC from hoopnet sampling in the LCR (GCMRC unpublished analyses). Second, a closed population mark-recapture experiment conducted in the LCR during the spring of 2001 indicated the population contained only 2,090 (95% C.I. 1611-2569; HBC >150 mm total length; USFWS unpublished data). Combined, these three independent analyses provide sufficient evidence to conclude that the Little Colorado River population of HBC is in decline.

Of paramount importance in conserving this population of federally endangered humpback chub is determining the factors contributing to this population decline and implementing management actions designed to minimize the effect of those factors. Although it is still unclear all of the factors that may be responsible for the recruitment decline beginning in 1992, we have identified a list of likely factors that could be acting either singly or in combination. These factors include: 1) Colorado and Little Colorado River hydrology, 2) infestation of juvenile HBC by asian tapeworm, 3) predation by or competition with warm-water native cyprinids and catastomids and nonnative cyprinids and ictalurids within the LCR, and 4) predation by or competition with cold-water non-native salmonids within the Colorado River.

The body of evidence available to evaluate specific hypotheses varies among the postulated factors. For instance, beginning in August 1991 the operation of Glen Canyon Dam was changed to reflect the so-called "interim operating criteria". This hydrology, and the subsequent Record of Decision flows that continue to present, can be generally characterized as having less severe daily flow fluctuations than the previous 28 years of load-following hydrology. Temporally, this major change in Colorado River hydrology correlates closely to the decline in HBC recruitment. Additionally, it is possible that the initial decline in HBC recruitment in 1992 was caused by the nearly continuous flooding in the LCR that occurred during summer 1992 through early winter 1993, particularly during the early summer time period when larval HBC emerge (Robinson et al. 1998). It is also possible that the high infestation rate of juvenile HBC by the introduced parasite asian tapeworm is a causative factor. HBC infected with asian tapeworm were first found during 1990, and infestation rates during 2001 have exceeded 90% (Anindo Choudury, pers. comm.). Finally, predation and competition by fishes either within the LCR or in the Colorado River may be driving the HBC recruitment trend. Although robust relative abundance data does not exist for non-native fishes within the LCR, there has been a large increase in the abundance of non-native salmonids in the Colorado River near the confluence of the LCR (LCR Inflow Reach 56.2 RM - 65.7 RM; Figure 2).

While it is difficult to determine which factor is most responsible for the HBC recruitment decline, a likely significant factor is negative interactions (predation and competition) with non-native fish. Interaction with non-native fish is implicated in the decline and extinction of native fishes throughout the Colorado River basin (Tyus and Saunders, III 2000 and references therein). Indeed, after being presented with the recent analyses describing the decline in the LCR HBC population, the Glen Canyon Dam Adaptive Management Work Group (AMWG) passed motions to begin planning and to conduct feasibility studies to reduce non-native fish abundance in the Little Colorado River and Bright Angel Creek. These first steps are commendable, however they do little to address the potential threat of predation and competition by rainbow (RBT) and brown trout (BNT) in Colorado River. To compliment these efforts, this study will be initiated to evaluate the potential effect of RBT and BNT predation on HBC recruitment and the efficacy of mechanical removal of RBT and BNT from the LCR Inflow reach.

Need

A series of experimental treatment scenarios for WY 2002-03 was developed by GCMRC in conjunction with the Adaptive Management Technical Work Group. At their April 24, 2002 meeting, the AMWG reviewed these scenarios and made their recommendation for implementing Experimental Flows and Mechanical Removal of salmonids in the LCR reach of the Colorado River Ecosystem. Secretary Norton approved the recommended experimental flows and mechanical removal in December 2002.

The recommended treatments contained within the experimental scenarios related to testing fish hypotheses center around the notion of improving future HBC recruitment by reducing the number of adult RBT and BNT residing in the system downstream of Lee's Ferry. Conceptually, this is to be accomplished in two ways. First, by reducing RBT and BNT recruitment by inflating the early life mortality rate of these fishes with highly fluctuating flows during their winter and spring spawning and rearing seasons. Second, by mechanically removing salmonids within the LCR inflow reach.

To date, a significant number of stakeholder groups have expressed concern about the winter and spring flow fluctuations called for in the experimental flows. Sport fishing interests are opposed to the fluctuating flows fearing significant negative impacts to the Lee's Ferry trout fishery. Additionally, several stakeholder groups have specifically asked: 1) whether or not reducing RBT and BNT abundance will improve HBC recruitment, and a related question 2) are RBT and BNT significant predators of HBC. This study is intended to address these questions as well as others formulated by the Technical Work Group (TWG) of the Glen Canyon Dam Adaptive Management Program. The TWG has identified a series of research information needs (RINs) specifically related to RBT and BNT predation on HBC. These include: "RIN 2.4.1-What are the most effective strategies and control methods to limit non-native fish predation and competition on native fish?; RIN 2.4.2-Determine if suppression of non-native predators and competitors increases native fish populations?; RIN 2.4.4-What are the target population levels, body size and age structure for non-native fish in the Colorado River ecosystem that limit their levels to those commensurate with the viability of native fish populations?; RIN 4.2.6-To what extent are RBT below the Paria River predators of native fish, primarily HBC? At what size do they become predators of native fish, especially HBC, i.e. how do the trophic interactions between RBT and native fish change with size of fish?" (GCMRC 2001). This work will attempt to answer some of these questions.

This study will also address a number of issues identified by the aquatic protocol evaluation panel (Anders et al. 2001). The panel had concerns with the lack of empirically established linkages between food base and fishes, and identified that a possible consequence of the recent increase in primary and secondary production may differentially benefit non-native species (competitors or predators) over native species. Secondly, the panel identified the need for establishing a better understanding for relationship and trophic linkages between foodbase and fish. Therefore, the trout dietary analysis in this study will be integrated with other existing GCMRC long-term monitoring programs that are the presently collecting or proposing to collect data specific to: 1) aquatic benthic foodbase, and 2) carbon productivity monitoring program.

Algae/macrophytes and invertebrates form the major components of the aquatic food base in the Colorado River ecosystem. The different macroinvertebrates consisting mostly of midge larvae (chironomids), black flies (simulids) and amphipods (Gamarus) trophically supports the trout fishery found in the Glen Canyon reach, as well as the fishery downstream of Lees Ferry. The foodbase is considered an important biotic resource because of the potential limitations, use, and availability required to support these different fish species. Research findings have revealed that a significant stair-step decrease occurs for both the composition and biomass of the major components of the foodbase (Usher and Blinn 1990, Hardwick et al. 1992, Blinn et al. 1993, Blinn et al. 1994; Shannon et al. 1994). This progressive decrease in the aquatic foodbase is related primarily to an increase in turbidity brought about from periodic tributary flows and the suspension of fines transported by higher discharges. Additionally, similar downstream patterns exist for fish distribution, compositional shifts, and a general reduction in relative abundance for certain species (Maddux et al. 1987, and Valdez and Ryel 1995). Separate studies have demonstrated a strong trophic linkage to the aquatic food base, as well as its spatial availability to fish (Shannon et al. 2001; Angradi 1994). Although, there is no direct evidence suggesting food limitations; studies measuring trophic pathways for the different biotic components have been conducted (Angradi 1994; Haden et al. 2002 In review; Shannon et al. 2001). These biotic patterns correspond to increasing distance downstream from Glen Canyon Dam.

Foraging preferences and nutritional requirements for these different fish species are not well known for this particular system (Anders et al. 2001). Certain observational studies (e.g., Valdez and Ryel 1995) have shown that

the overall assemblage of fish use different aquatic and terrestrial invertebrates, as well as fish that are either young or small-sized (Valdez and Ryel 1995; Rowell 2001). It is ecologically recognized that most young developing fish do not survive to recruit into the reproductive population. Most of the mortality occurring to these vulnerable fish is due to predation. Therefore, small sized fish represent a proportion of the overall foodbase in this ecosystem. The physical and biotic factors that regulate their availability as a food item, as well as their survival, influence the population dynamics of these different fish species. Although predation has been documented for the different trout species (rainbow and brown trout), their apparent food habits as indicated by stomach content analysis are not conclusive. Especially, when it comes to understanding the possible trophic interactions that exist between different size-age classes and the environmental pressures associated with different population densities and variable food availability.

Our current understanding regarding benthic and drift production and fish life histories limits our ability to make these linkages between lower trophic levels and food availability for native and non-native fishes (Anders et al. 2001). Although it has been implicitly recognized that fish consume invertebrates and fish, previous research has not demonstrated food limitations to higher trophic levels (Valdez and Ryel 1995). Because the assumed trophic linkages between food base and fishes have not been empirically established, it is difficult to determine food base requirements (Anders et al. 2001). Therefore, recent increase in algae/macrophytes and invertebrates may have direct benefits to native fish. Conversely, the process of maintaining or maximizing the production of the aquatic food base may benefit solely non-native species that are possibly better competitors, and/or predators in this altered ecosystem.

OBJECTIVES

The study is motivated by the following classes of objectives: 1) Efficacy of mechanical removal of adult RBT and BNT from the LCR Inflow reach, 2) RBT and BNT predation, and 3) Effect of adult RBT and BNT in the LCR inflow reach on the population dynamics of the LCR HBC population.

Efficacy of Mechanical Removal of Adult RBT and BNT from the LCR Inflow Reach

- 1. Estimate abundance of adult RBT and BNT in the LCR Inflow reach prior to each removal event.
- 2. Estimate changes in adult RBT and BNT size composition in response to removal events.
- 3. Determine trout immigration rate (Seasonal and Annual) into the LCR Inflow reach between removal events.
- 4. Estimate gear efficiency as a function of boat type, turbidity, season, and dominant habitat type.

Rainbow and Brown Trout Diet Analysis and Predation

- 1. Estimate the instantaneous proportion of adult RBT and BNT residing in the LCR Inflow reach that are piscivorous.
- 2. Determine relationship between adult RBT and BNT total length and likelihood of piscivory.
- 3. Estimate the relationship between adult RBT and BNT total length and gape.
- 4. Estimate the relationship between adult RBT and BNT total length and prey body depth.
- 5. Estimate adult RBT and BNT diet composition.

Effect of Adult RBT and BNT in the LCR Inflow Reach on the Population Dynamics of the LCR HBC Population

- 1. Evaluate the relationship between adult RBT and BNT abundance in the LCR inflow reach and juvenile HBC survival/retention rate in the LCR inflow reach.
- 2. Evaluate the relationship between adult RBT and BNT abundance in the LCR inflow reach and recruitment to the LCR HBC population.

METHODS

Study 1. Procedures for the Mechanical Removal of Non-Native Salmonids from the Little Colorado River Reach of the Colorado River

Study Area and Design

Removal Reach: The LCR Inflow reach is recognized for having the highest abundance of adult and juvenile HBC in the Colorado River mainstem (Valdez and Ryel 1995). We have selected a study area (56.2 RM - 65.7 RM; Figure 3) that encloses the majority of the geographic distribution of the LCR HBC population. The study area is stratified into 6 river reaches: A-F. Reaches A and B are the right and left shore reaches from Kwagunt Rapid (RM 56.2) to Science Beach (RM 61.5). Reaches C and D are the right and left shore river reaches between RM 61.5 to RM 62.1 and include the LCR confluence and the mixing zone below the LCR. Reaches E and F are the right and left shore reaches downstream of the LCR confluence (RM 62.1 to Lava Chuar Rapid RM 65.7). We stratified the study area into these 6 reaches in order to control for the affect of the LCR discharge into the Mainstem Colorado River. Reaches A and B are unaffected by the tributary and reaches E and F are believed to be of sufficient distance downstream of the mixing zone to be affected uniformly throughout. Reaches C and D include the LCR confluence and will be differentially affected by LCR discharge throughout their lengths. Within river reaches A-B and E-F, the shoreline is divided into 500m sites. The number of sites within each river reach is as follows: A=19, B=19, E=13, and F=14 (13 shoreline sites and one island site). Reaches C and D constitute single sites. A base camp for each trip will be established at Science Beach (across from the LCR confluence).

The upstream and downstream study area endpoints are bounded by hydraulic and geomorphic control; however, they are not impermeable to system-wide fish movement (Stevens et al. 1997). For this reason, depletion efforts will be conducted that are both spatially discrete, and repeated seasonally over a period of 4 years (Table 1). We will conduct annually, three depletion trips in January-March and three depletion trips in July-September. The annual depletion efforts will be repeated four years, for a total of 24 times, to determine how removal of fish using a series of depletion passes in a discrete area will influence the relative abundance of the remaining fish stock. Since we will be unable to control for migration, recruitment and mortality occurring at a local level, comparisons among trip population estimates will be analyzed in order to evaluate if mechanical removal methods are an effective means to control for undesirable fish species. The sampling efforts are scheduled to coincide with the major periods of LCR flooding events (spring runoff and monsoonal storms) that are correlated with juvenile HBC immigration to the mainstem Colorado River (Valdez and Ryel 1995).

Control Reach: To determine if differences in fish population characteristics (e.g., relative abundance, size structure, etc.) in the experimental reach is a function of environmental influences/fluctuating flow treatments and not the mechanical removal, a control area has been selected (44 RM – 52 RM; Figure 4) and divided into 60 500 m sites occurring on both sides of the river. The 24 randomly selected sites within the control area will be sampled to estimate the relative abundance and size structure on the first night of each trip (once per trip, 6 times per year). All fish collection, handling procedures, and data recording will proceed as described below for the removal reach except no fish will be euthanized within the control reach.

Data Collection

Control Reach: Table 2 details the day specific tasks to be performed within a typical trip. On day one, sport boats will proceed ahead of the group to distinguish and mark control sites to be sampled. On night one, electrofishing within the control reach will begin. Four boats will electrofish a total of 24 sites (2 boats 7 sites each and 2 boats 5

sites each; Table 3a) to complete all data collection in the control reach. The transport boat will assist those boats sampling 7 sites with fish processing in order to speed overall sampling activities. All fish will be measured to determine relative abundance and size structure within the control reach. All fish collection, handling procedures, and data recording will proceed as described below (for the removal reach) except no fish will be euthanized within the control reach. Additionally, each captured rainbow and brown trout greater than or equal to 200 mm will be fitted with a floy tag between the dorsal fin pterygiophores near the posterior portion of the fin. The tag number and recapture status will be recorded in the two fields associated with Tag2 on the Electrofishing Depletion Data Sheets (Appendix A). All RBT and BNT fitted with a flow tag will also be given a left Pelvic Fin Clip (LP2). Finally, trout processed by the transport boat crew will be weighed in order to collect the data necessary to estimate RBT condition in the control reach. There will be no processing station; upon capture, all fish will be placed in fresh water to be worked up at the end of each sampling site. Each electrofishing boat driver/netter will be responsible for their data collection and recording on the Standardized Depletion Electrofishing Data Sheet (Appendix A). When each site is completed, all fish will be released at the upper end of the site.

Removal Reach: Following arrival at the Science Beach base camp on day two, GPS units and aerial photographs will be used to mark the boundaries of the 500m sites within reaches A-B and E-F (Figure 3). The boundaries will be marked by hanging lengths of pvc pipe wrapped with reflective tape. Each of these boundaries will also be marked with small aluminum tags inscribed with the site name immediately downstream of the boundary. Over the course of 10 nights (Days 3-12 in Table 2), all sample units within reaches A-B, E-F, and C will be electrofished 5 times. The upper ½ of reaches A-B and E-F and reach C will be electrofished on days 3,5,7,9,and 11. The lower ½ of reaches A-B and E-F will be electrofished on days 4,6,8,10, and 12. Reach D will not be electrofished unless low water conductivity and low native fish abundance can be assured. Electrofishing is not to begin earlier than dusk. The actual start time will depend on the sampling season and will be decided upon at the beginning of each trip. Electrofishing boat operators are to ensure that boats are maintained and fully operational prior to electrofishing. On each of the four (4) boats all of the necessary electrofishing equipment and supplies are to be checked prior to use. This is the responsibility of the technical electrofishing boat operator. Upon completion of nightly electrofishing, the electrofishing crews are to transport the remaining catch to the Processing Station.

Boat, Driver, and Netter Allocation within Reaches: A total of 4 electrofishing boats of two types each will be utilized in this study. The boat types are: 1) 15' Achilles sport boat (rubber hull) and 2) 15' Osprey sport boat (aluminum hull). Within the removal reach, a rubber boat and an aluminum boat will always be used above the LCR and 1 of each will always be used below the LCR. The 4 boat drivers will be randomly assigned to a particular reach/depletion run within each trip. The underlying purpose for the random assignment is to control for systematic bias that might exist among different electrofishing boat operators. Electrofishing boat operators are expected to sample all 10 nights without substitution or assistance from other boat operators on the trip (emergencies or illness aside). Table 3b reports boat type and boat driver assignments by depletion run for the January, February, and March trips. This design will assure that accurate assessments of reach and boat specific catch-rates are not biased by (or can be adjusted for) driver and netter affects. Sampling equipment, methods and electrical configuration used will be consistent with the established GCMRC fish handling and sampling protocols (Appendix E).

Netting is an extremely important component to the success or failure of this study; therefore this activity requires attentive behavior. However, no attempt will be made to control for variability that exists among netters. Only a single netter will be used per boat. Technical personnel that are netting are expected to rotate out every other day through the different fish electrofishing and processing activities. A personnel schedule will be posted at the science camp identifying a rotation schedule. Netters are expected to rotate between different boats, electrofishing boat operator and processing activities (refer to posted work schedule). Netters are expected to be consistent in their performance, which requires being safe, observant and coordinated. Owing to the continuous workload, if a netter becomes fatigued, alternates will replace he/she during the night. The electrofishing boat operator will make this decision. The transport-boat will be used for exchanging netting personnel rather than the electrofishing boat.

Electrofishing Power Standardization: The two boats will be outfitted with identical Coeffelt CPS mark XXII electrofishing boxes. Since the boats produce different electrical field characteristics due to different electrode configurations and therefore fishing efficiencies, the same boat will be used for each depletion run within a river reach for an entire trip. In an attempt to standardize efficiency as much as possible, the power output for each boat will be adjusted to produce on average 5,000 watts (e.g. Achilles 333 volts, 15 amps; Osprey 238 volts, 21 amps). technical boat operators will be supplied with a power curve to allow them to standardize power to 5000 watts for any ambient water conductivity (Figure 6). Power should be standardized in mid-river or a suitably deep location so

that the electrical field is free of obstructions during standardizing. Power will be standardized when the CPS unit has been adjusted so that values for voltage and current (amperage) fall on the 5000 W power curve.

Fish Handling Procedures: During removal operations, qualified personnel are to identify fish to species. A fish key will be available for reference. Fish are then to be separated into two storage containers as either native- or nonnative fish. Salmonids and other non-native fish (catfish, carp, fathead minnows, killifish, etc.) are to be euthanized and temporarily stored in 1/8" small mesh net. These small-meshed Net Sample Bags (30-L capacity) are to line the inside of the non-native fish storage container containing the euthanizing solution. Since there is a potential for inadvertently placing native fish in the euthanizing bath, netters are to collect no more than two fish per net sweep to avoid potential misidentification of native fish. Upon completing the designated electrofishing site the smallmeshed Net Sample Bags are to be removed and transferred to the shoreline above the high water mark at designated collection points. A Data Info Card with all pertinent information will be filled out and attached to the sample bag containing euthanized fish. These data are to include: Study Reach (A, B, C, D, E, or F) and section # (1-19), date, depletion #, and total effort (seconds). In addition to the Data Info Card, a light stick will also be affixed to the bag. If an electrofishing run did not catch fish, an empty net and accompanying information will be left at the designated collection point for the fish transport boat. Additionally, if an electrofishing run did not catch fish, a depletion electrofishing data form will also be completed to record sample specific information (e.g. site, effort, depletion #, etc.). This is to be done to avoid any undue confusion regarding missed sites. The locations for these fish collection points are to be at or near the end of the electrofishing section and are to be visually apparent (i.e. light sticks visible from the river). Sample bags are to be rocked down so as to avoid scavengers (coyotes and foxes) inadvertently displacing fish carcasses. This is the responsibility of the technical electrofishing boat operator.

Non-native fish are to be euthanized using a solution super-saturated with carbon dioxide (CO₂). The solution will be prepared by dissolving 6-8 oz of sodium bisulfate (NaHSO₄; swimming pool acid) in approximately 12 gal of water to make a slightly acidified solution. Two pounds of sodium bicarbonate (NaHCO₃; Baking Soda) contained in a perforated zip lock bag will then be sunk to the bottom of the container. As the acidified water reacts with the baking soda, the solution will effervesce, driving the oxygen out of the water and causing the solution to become super-saturated with carbon dioxide. This solution should be effective for an entire night's work, but may require freshening with additional baking soda if used for multiple days.

Native fish caught during the electrofishing run are to be separated and placed in a separate container containing fresh water. Native fish will be processed and released alive in the field by qualified boat personnel. Standard fishery measurements are to be collected on all native fish encountered. To avoid recapture, fish will be transported to the upper extent of the electrofished section. Passive integrated transponders (PIT) are used as a method for mark-recapture estimates for abundance, relative year class strength, recruitment, growth and movement in the Colorado River mainstem and associated tributaries. All native fish (>120 mm) will be assessed for PIT-tags; as well as fin-clips and other associated marks being used as part of the GCMRC monitoring program. To avoid depressed oxygen levels, water needs to be periodically changed. This should occur between individual electrofishing runs. In the unlikely event of native fish mortality (e.g., endangered humpback chub), the specimen(s) will be preserved and brought back to the GCMRC for a complete analysis. Specimen(s) will be documented in the field and standard measurements (PIT-tag, TL (mm), weight (g), sex, and the presence of external parasites) will be collected. A visual inspection will be conducted to evaluate the fish for any skeletal abnormalities, or bruising and discoloration that may occur from electrofishing. A necropsy will be performed to assess for abdominal and intramuscular injuries. The fish will be eviscerated carefully removing stomach and entire intestine for later dietary analysis.

Data Recording: Boat drivers/netters will be responsible for filling out Electrofishing Depletion Data Sheets following electrofishing each site even if the site produced no fish to record sample specific information (e.g. site, effort, depletion #, etc.; Appendix A). Data sheets are to be legible and completed for each electrofishing section. In order to expedite electrofishing effort these data sheets have been simplified to avoid redundancy in data recording. Data sheets are to remain on each of the four-electrofishing boats between runs. Upon completion of the nightly electrofishing activities, each of the electrofishing boat operators are responsible for transferring all completed data sheets to the data storage box. The following day the electrofishing boat operator/data recorder is to evaluate all recorded data sheets and identify any errors that might have occurred during the previous nights transcription. Upon validation, the data sheets are to be placed in the designated Data Storage Box for safekeeping. Any corrections made to the data sheets need to be drawn to the attention of the designated Trip Principal Investigator (PI). It is the responsibility of the trip PI to ensure that all data sheets have been properly filled out by electrofishing boat operator and processing crew.

Fish Transport: A transport boat is to collect fish at each of the designated collection points; however, because of the distance between collection points the boat operator will alternate between the upper and lower reaches while transporting fish to the Fish Processing Station. All attempts will be made to avoid undue gas consumption. The return time interval between sampled reaches is estimated at 1 to 1.5-hr. All non-native fish will be transported to Processing Station for data collection activities and processed during a single electrofishing night to avoid decomposition and loss to stomach samples. Transport boat personnel are to verify if the Data Info Card with all pertinent information has been filled out and attached to the sample bag containing euthanized fish. These data are to include: Study Reach (A, B, C, D, E, or F), site # (1-19), date, depletion #, and total effort (seconds). If data is absent this information must be acquired from the responsible electrofishing crew. Additional responsibilities include the collection of drift during nights 1 & 2. The availability of drift is to correspond with the first depletion effort. Boat operator will alternate drift sampling between upstream and downstream reaches at 12 established sites. A total of six sites, three upstream and three downstream are to be sampled per night, and then repeated at six other sites the following night. Each of the sampling locations is marked by a set of buoys that are sufficiently anchored within the mid-channel eddy complex. Buoys are to provide for a secure boat attachment and to be used as a stationary sampling platform. A total of six replicate samples are to be collected over the course of a 6-h period. This drift sampling is to coincide with the transport of euthanized fish to the Fish Processing Station. This sampling effort requires two personnel in order to deploy multiple samplers, collect and data record. Samples are to be taken to Fish Processing Station for sieving and preservation (refer to Drift Sampling). The drift effort is to be discontinued after the second night.

US Fish and Wildlife Fish Handling Directives: As a condition of the Biological Opinion (USFWS 2002) associated with mechanical removal operations issued by the US Fish and Wildlife Service, the following procedures must be followed at all times:

- 1. All humpback chub captured will be held separately from non-native fishes to minimize stress, predation, and injury during recovery from electrofishing. If this cannot be accomplished, the non-natives shall be sacrificed.
- 2. In the Control Reach, action agencies shall provide the greatest release distance between native and non-native fishes as possible.
- 3. All humpback chub shall be processed and released immediately after recovery in the near-shore habitat where they were collected.
- 4. All rainbow trout captured in hoopnets shall be checked for predation on humpback chub.
- 5. Placement of pit tags by field crews shall be conducted only by those individuals previously permitted by the Fish and Wildlife Service to handle and pit tag humpback chub, or those individuals who have received training by permitted individuals and 20 hours of supervised pit tagging.

Data Analysis

A variety of data analyses will be performed to address the objectives under: efficacy of mechanical removal of adult RBT and BNT from the LCR Inflow Reach. The major data analysis tasks and associated strategies include, but are not limited to, the following:

Task 1. Estimate initial abundance in each of 4 river reaches (A-B and E-F) for each trip during 2002 – 2005. We will conduct five-pass Leslie DeLury depletion estimates for each site within each reach (Van Den Avyle and Hayward 1999). The mean population estimate (number of trout per 500 m of shoreline) will be calculated for each reach and compared among reaches and years with an analysis of variance (ANOVA).

Task 2. Test whether initial abundance varies between depletion trips. The mean population estimate (number of trout per 500 m of shoreline) will be calculated for each reach from each individual site population estimate (n=13-19, depending on reach) and compared among reaches and trips with a two-way repeated measures analysis of variance (ANOVA) with reach and month as the class variables. Because each site within each reach will be

sampled more than once, the population estimate from each site may not be independent of the population estimate from that same site subsequent sampling trips. Therefore, sampling site will be the repeated variable.

- Task 3. Evaluate differences in efficiency among boats (rubber versus aluminum). The catchability coefficient (q) will be calculated for each 5-pass depletion estimate at each site within each reach. A three-way ANOVA with interaction terms will be used with boat type, habitat type, and river reach as main effects and q as the dependent variable. A significant (P<0.10) boat effect would suggest that boat type influenced catchability (assuming there were no significant interactions that included boat type).
- Task 4. Evaluate differences among initial abundance among dominant shoreline habitat types. The shoreline of the entire study area will be classified into four discrete habitat types based on existing geographic information systems models. Each sampling site will be then classified as the dominant habitat type found in that site. If there is not a dominant habitat type (e.g., equal length of two habitat type within the same site), this site will be removed from further analysis. A three-way ANOVA (with interactions) will be used to compare depletion population estimates among habitat types, two river reaches (upstream of LCR and downstream of LCR), and sampling trip.
- **Task 5.** Evaluate differences in efficiency among turbid and clear water conditions. Catchability (q) will be calculated for each depletion estimate at each site within each reach. An analysis of covariance will be used to determine if q decreased with increased turbidity, controlling for river reach. In this analysis, we hypothesize that q will increase with increased turbidity. However, q may also be affected by river reach; therefore, river reach will be used as a covariable.

Study 2. Salmonid Diet Analysis and Invertebrate Drift

There are two separate components associated with this salmonid diet/predation data collection effort; they are: Total Trout Diet Analysis and Trout Piscivory Analysis. Methods to achieve these analyses are described below.

Fish Processing and Data Collection

As described above a fish processing station will be set up at the science camp (61.4 RM). At the processing station one individual will function as field coordinator to oversee all processing activities. A four (4)-person crew will be responsible for all data collection and fish processing activities. These personnel include 1) data recorder, 2) length/weight measurements, 3) stomach evisceration/preservation, and 4) sample organizer. All processing activities will occur during night and are estimated to extend over an 8-hr period.

Data Recorder

This individual will be in charge of organizing supplies and equipment, replenishing of stock supplies, personnel scheduling, and data recording. The designated processing personnel are to assist in all aspects of these field tasks. All data will be recorded during each processing session/night. The following day the data recorder will check all recorded data sheets to identify any errors that may have occurred. Upon finalization data sheets are to be placed in safekeeping in the designated storage box. Trip leaders are responsible for examining data sheets following completion and verifying that they have been filled out properly.

Measurements: Specific measurements on all sampled fish will be recorded at the processing station. These measurements include: numeric-coded tag for stomach sample identification, species identification, total length (mm), weight (g), sex, and gape-width (mm). Information on recaptured fish such as PIT-tagged trout and fin clipped trout will also be recorded. All measurements taken will be called out to the data recorder and entered into the Fish Processing Data Sheets (Appendix B).

Stomach collection: Stomachs from all non-native fish will be collected, preserved, and stored in nalgene containers to ensure that no contents are lost. The preservative solution will consist of 95% ethanol prepared in advance. It has been estimated that 25-ml of preservative will be used for each collected stomach; however, adjustments may be made according to size and contents. A length-wise incision will be made along the foregut to ensure preservation. Each stomach will have a corresponding numeric metal-coded tag placed in the bottle with all contents. Fish will be processed sequentially and assigned to a sampling reach and section, referred to as a sampling lot. All fish collected during an electrofishing run will be collectively separated and stored using a larger plastic

bag. Each plastic bag will be marked according to the designated sampling reach, site and date. Aluminum storage boxes will be used as storage containers for the different Sampling Lots. These storage boxes contain 380 nalgene containers that have been pre-filled with ETOH preservative. Each storage box will have an information sheet that identifies all Sampling Lots contained. Information will include Sampling Lot number and sampling date. These sampling lot data sheets will be stored in a designated folder in the data box. The numerical metal-coded tag assigned to each fish/stomach will allow identification back in the lab. All data recorded at the processing station for a specific fish will be linked to the stomach contents via the metal numeric-coded tag. Stomach samples used for diet analysis are to be representative of normal feeding behavior. Yet, it is conceivable that some of the fish during a depletion trip may have been exposed to multiple depletion passes before actual capture due to gear and netting This becomes problematic since feeding behavior may be disrupted by multiple exposures to electrofishing. Therefore, stomach samples used for diet analysis are to be collected only from the first depletion pass (night 1 & 2) for a given trip. To avoid assessing digested material associated with the lower intestine, only stomach contents found in the foregut are to be used for diet analysis. Therefore to discriminate between foregut and intestine, gut contents are to be separated, stored and preserved in different nalgene containers. Intestinal tract is to be separated at junction of foregut and intestine using scissors, and then each region is to be incised lengthwise and separately placed in designated containers. Initial step requires placement of a metal numeric tag face down in bottom of 125 ml container followed by incised foregut. Secondly, the smaller intestine is also to be incised and placed in a 60 ml container and sealed. The smaller container is to be inserted into the larger 125 ml container and sealed. All containers have been pre-filled with ETOH prior to trip. The purpose for this is to maintain the association of the different stomach contents to the same numbered tag. For redundancy, processors are then to mark the outside of the larger nalgene container using the same numeric tag number. This double mark is only necessary for gut contents used for diet analysis. Alternately, the entire intestinal tract is to be used for assessing predation, therefore, fish captured during the remaining depletion passes from night 5 to 12 are to retain the entire intestinal tract. This will be incised, stored, and preserved as described above.

Fish Disposal

Fish carcasses will be temporarily stored in sealed garbage containers after each processing night. The following day fish carcasses will be mechanically ground and placed in fish disposal barrels (15 gallon). A Honda 5,000 w generator will be used to power the electric Weston grinder (Model 32, 115v, 1.5-hp). The disposal area will be located at the downstream inlet at Science Beach (61.4 RM). A sturdy aluminum table (20"x 44") and drop cloth will be used for the fish grinding and disposal process. Only personnel trained and familiar with its safe operation are to use the grinder. Ground-up carcasses will be fixed using phosphoric acid to prevent decomposition and later storage problems. These plastic containers are to be stored on a designated boat. Polyethylene tubing is to be used as a manifold to interconnect disposal barrels. Excess tubing will be extended into the river to bleed off any excess methane gas build-up. Owing to caustic nature of concentrated phosphoric acid only GCMRC will be responsible for dispensing of preservative. The disposal area will be kept clean so as to avoid undesirable health issues and attracting scavengers. Liquid Clorox bleach will be used for washing and disinfecting garbage cans and the processing area. All excess material will be collected, washed and disposed of using available brushes and bleach. Containers will be stored and transported from the Grand Canyon after each mechanical removal trip. All material produced from grinding will be delivered to the Hualapai Nation for fertilizer following the trip.

Upon trip completion the USFWS will be notified of the incidental take as specified in the research permit. Additionally, these fish will be assessed for skeletal abnormalities using radiogrammetry techniques (Sharber and Carothers 1988; Sharber et al. 1994). Specimens are to be sent to Arizona Game and Fish Department for the purpose of cataloguing and transferring such material to a proper facility for permanent collection and curatorship.

Trout diet analysis

Certain research questions exist regarding trout diet differences among species, age-size class structure and location. This type of data can be very useful in developing an energetics model. Preserved fish stomach contents will be used for this assessment using a stratified random sampling approach, whereby stomach samples are to be stratified by species (RBT and BNT), reaches (Above-LCR and Below-LCR) and length groups (< 150 mm, 151 - 200 mm, 201 - 300 mm, and > 301 mm; Table 4). Following the completion of the trip, 240 samples are to be randomly selected from the total number of samples collected during night 1 & 2, for the purpose of separating, categorizing, and enumerating identified items. These categorized items are to be desiccated for 24-h at 60°C, weighed (± 0.1 mg), ashed for 1-h at 500°C, and reweighed for ash-free standing mass determination. Digested fish and bones

collected from stomach contents of non-native species will be used as voucher and diagnostic material. And where possible such material will be used for taxonomic purposes.

Laboratory Procedures: Gut contents will be analyzed from a set of sub-samples that are randomly selected and stratified by sampling locality, time and fish size (Table 4). Sample size is designated by fish TL (mm) and study reach. The diet analysis will quantify all ingested phytobenthic material, macroinvertebrates, and vertebrates using a combination of analytical methods (volumetric, weight, and numeric counts) taxonomically identified (Marrero and Lopez-Rojas 1995; Rowell 2001).

Seasonal and inter-annual differences in the availability of the aquatic food base (standing biomass and drift) are to be linked to fish feeding habits and electivity preferences. Additionally, stomach samples will be collected in the Lees Ferry Reach to assess diet. The Lees Ferry Reach is to serve as a spatial control lacking effects from the fish removal occurring downstream. The collection of these stomach samples will be coordinated with the Lees Ferry monitoring program, conducted by Arizona Game and Fish, Department.

The metal numeric-coded tags assigned to each stomach sample will be linked to the data collection effort and used for stratifying and randomizing the samples selected for diet analysis. If stomachs are observed to be empty an additional sample will be resampled from the underrepresented strata.

Sampling problems may occur. It is understood that BNT abundances are much lower than RBT in the LCR-inflow and may be underrepresented. Additionally age-class structure will be skewed toward larger fish that may have migrated into the region from the Granite Gorge.

Detailed stomach analysis will be achieved for 30 trout per length group for each reach and species; however, all collected stomachs will be analyzed for the presence or absence of fish or fish remains. Special dye markers (Alizarin red and KOH) will be used to highlight bones and cartilage contained within the gut contents. The effectiveness of the stain is rate dependent, so that at the concentrations (0.1%) used, the preserved material should be allowed to fix over a 24-hr period. Where possible, bones will be used for reconstructing and identifying prey taxa.

All collected specimens and data sheets are to be assessed for completion, accuracy, and data entry errors, and sample specimens are to be cataloged, organized and stored for later transport. All data will be entered following trips consistent with GCMRC format structures.

Data Analysis

Statistical analyses are to be performed to address the objectives under rainbow and brown trout diet analysis and predation. Based on previous research, age-0 HBC abundance will likely be higher below the LCR than above the LCR. Therefore, we hypothesize that the incidence of HBC predation will be higher below the LCR. However, other factors that may influence HBC predation may be trout species (rainbow or brown), trout length, trout and HBC abundance, and turbidity. Therefore, a logistic regression model (Agresti 1990) will be used with the dependent variable being presence or absence of HBC in a trout stomach. This model will include the variable suggested as potential factors that may influence HBC predation as listed above. Comparisons made among seasons and within years will provide information on whether or not particular cohorts are more vulnerable to predation due to differences in size, relative prey abundance or relative predator abundance.

Drift sampling

The drift sampling effort is to provide a representative sample of the available drift for foraging fish. Sampling will be conducted by the transport boat so as not to interfere with electrofishing duties. Sampling will be conducted at 12 sites (6 above the LCR and 6 below the LCR). At each of these sites, four replicate samples will be collected with a 30-cm x 120-cm width to length (1:4 ratio) and 363 µm mesh size plankton net with a current meter attached near the center of the net opening. Drift nets are to be inserted into an aluminum frame for ease of deployment. These nets will be deployed in eddies directly adjacent to the electrofishing stations using a lateral hinged boom off the sides of the boat between MT 1700 and 2200. Samples are to be collected over a 5-min period at flow velocities of 0.15-0.25 m s⁻¹. Samples will be collected 2 times (nights 1 & 2) during the non-native depletion sampling for a total of 48 samples during each trip (Table 5). Data collection will include: station location, date, time, flow velocity,

depth, drift net number. In the field, drift contents are to be initially containerized using the drift collection bucket and sealed. These will be transferred to the processing station for sieving (363 µm mesh size) and preservation in 10% ethanol.

Study 3. Procedures for Estimating the Relative Abundance of Juvenile HBC in the LCR Inflow Reach

Mini-hoopnets will be used to estimate the relative abundance of humpback chub at the 30 standardized sites downstream of the LCR confluence. Data obtained during past investigations suggest that relative abundance estimated at these sites over several months (September, November, and January) may be useful in estimating the mortality/emigration rate of juvenile HBC from the LCR inflow reach (Valdez and Ryel 1995). In addition, these collections, along with incidental catch of juvenile HBC during electrofishing, will provide size structure information for juvenile HBC in the LCR inflow reach.

Study Area and Design

30 hoopnets will deployed for 3-24 hour sets on the 1st, 3rd, and 5th days of the trip. Set locations will correspond to the 30 standardized locations established by Gorman and Coggins (2000; Figure 5). Deploying nets on the 2nd, 4th, and 6th days of the trip will allow the nets to be fished during time periods when electrofishing activities are not being conducted within the sites that are occupied by the nets. The nets will be deployed between 1100 and 1300 and retrieved the following day (i.e. trip days 3, 5, and 7) during the same timeframe.

Data Collection

Results from the hoopnet sampling will be recorded on the standard netting data form (Appendix C). All captured fish will be handled and processed according to the procedures detailed in Ward 2002 (Appendix D). Upon completion of the hoopnetting activities, the boat operators are to be responsible for transferring all completed data sheets to the data storage box. Before the next netting effort is initiated, boat operator/data recorder is to evaluate all completed data sheets and identify any errors that might have occurred during the previous efforts transcription. Upon validation, the data sheets are to be placed in the designated Data Storage Box for safekeeping. Any corrections made to the data sheets need to be drawn to the attention of the designated Trip Principal Investigator (PI). It is the responsibility of the trip PI to ensure that all data sheets have been properly filled out.

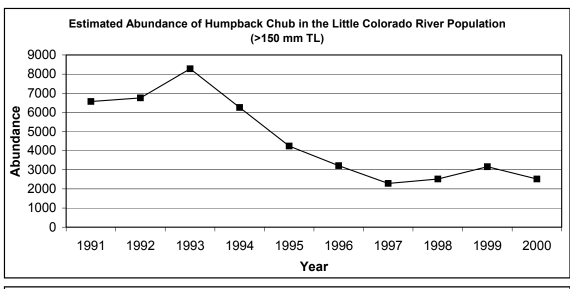
SCHEDULES AND REPORTS

Semi-annual reports and presentations will be given to the AMWG and/or TWG during December and June. At least 4 publications in the primary literature are expected as a result of this work to be submitted beginning in 2004.

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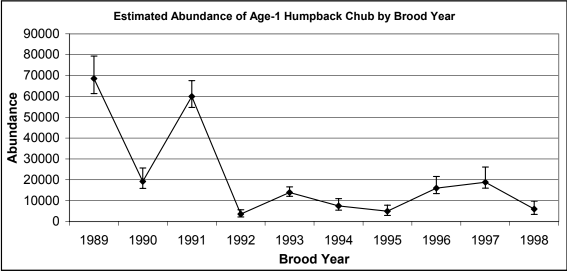


Figure 1. Estimated annual trend in population size (top panel) and recruitment (bottom panel) of the Little Colorado River population of humpback chub, Grand Canyon, Arizona.

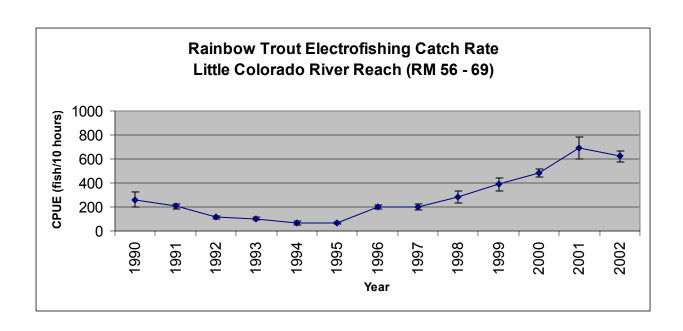


Figure 2. Relative abundance (number of fish per 10 hours of nighttime electrofishing) of rainbow trout (top panel) and brown trout (bottom panel) in the area of the Colorado River near the Little Colorado River confluence, Grand Canyon, Arizona, 1990-2002.

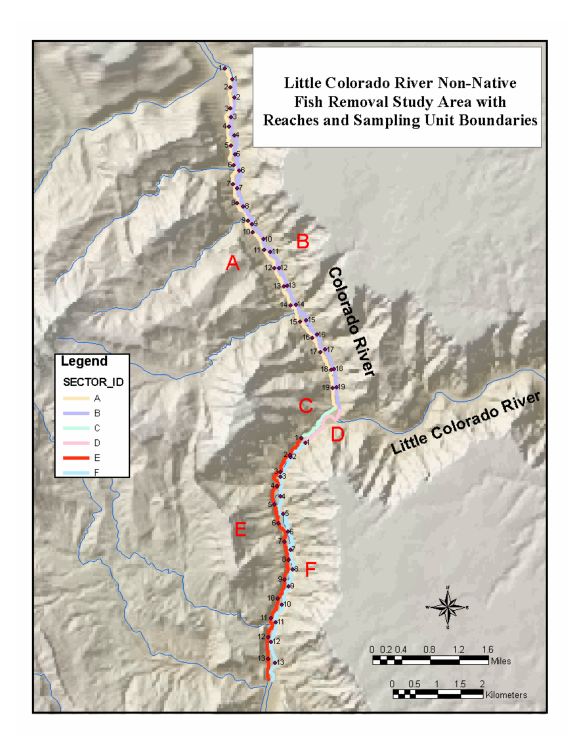


Figure 3. Proposed study area for the mechanical removal of non-native fishes in the Colorado River near the confluence of the Little Colorado River, Grand Canyon, Arizona. Six study reaches are delineated (A-F) and each sites within each reach is 500 m.

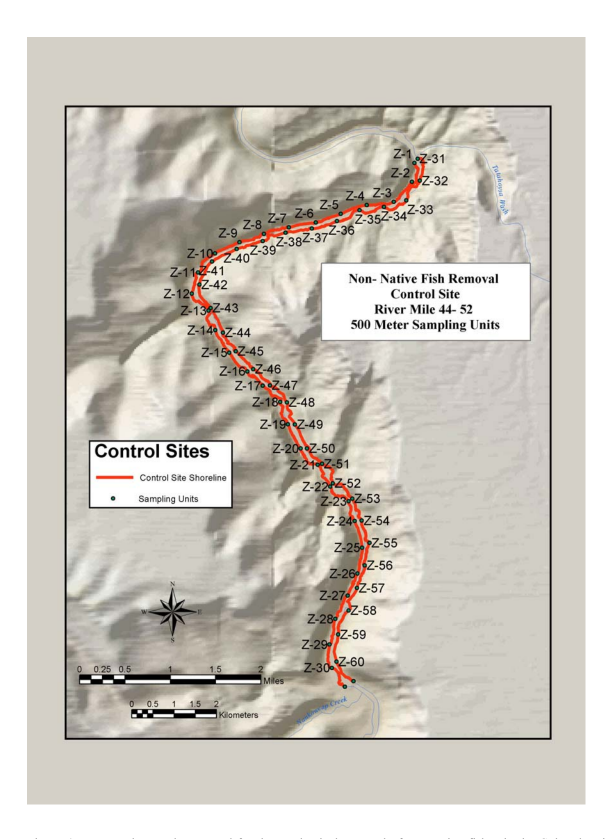


Figure 4. Proposed control area used for the mechanical removal of non-native fishes in the Colorado River, RM 44-52, Grand Canyon, Arizona. Randomly selected 500 m sites will be electrofished from this area each sampling trip; however, no fish will be removed.

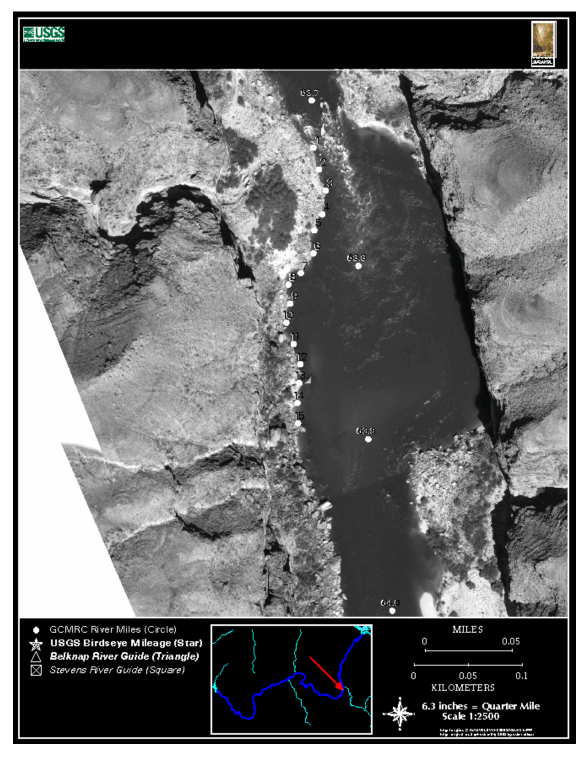


Figure 5a. Hoop-net locations (stations 1-15) for standardized humpback chub sampling in the Colorado River below the Little Colorado River confluence, Grand Canyon, Arizona.

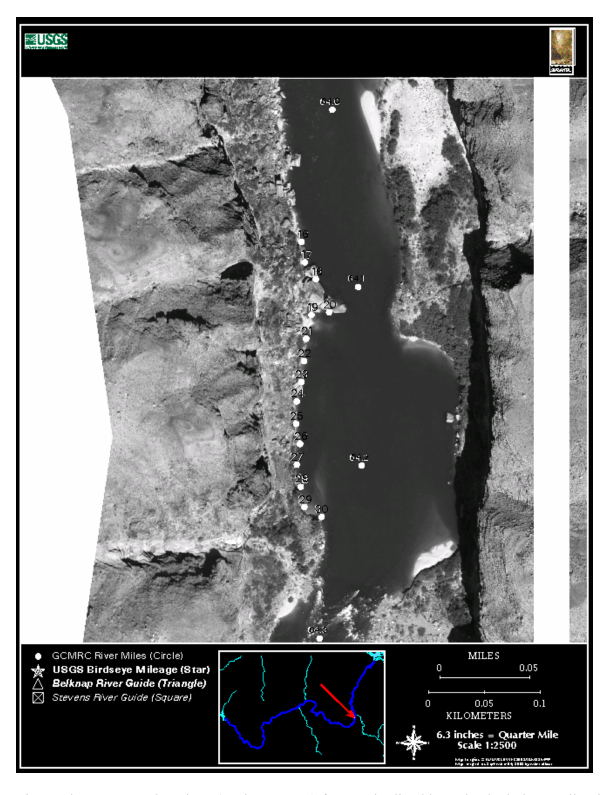


Figure 5b. Hoop-net locations (stations 16-30) for standardized humpback chub sampling in the Colorado River below the Little Colorado River confluence, Grand Canyon, Arizona.

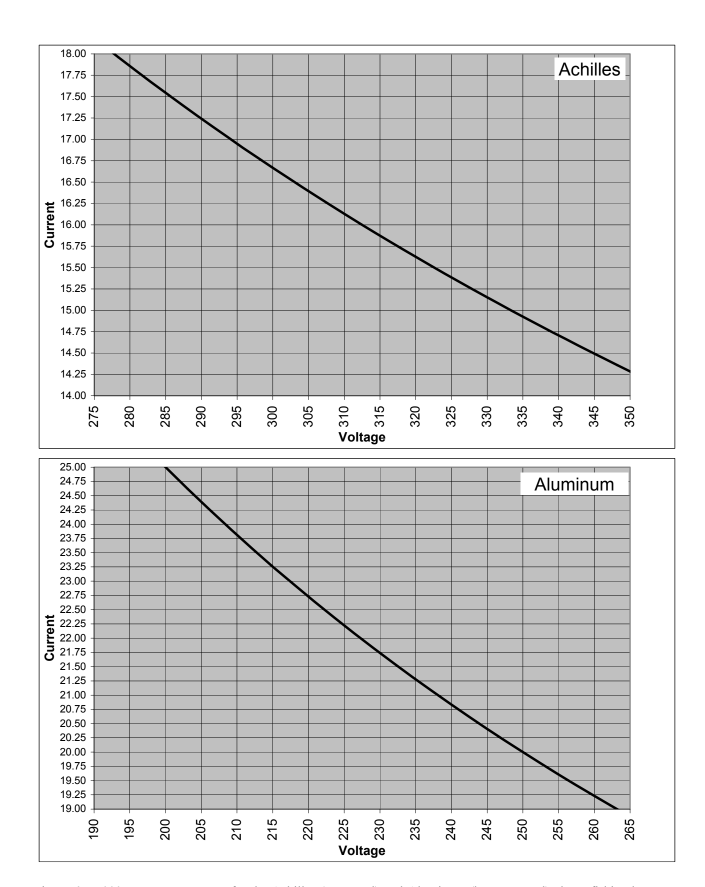


Figure 6. 5,000 watt power curves for the Achilles (top panel) and Aluminum (bottom panel) electrofishing boat. CPS settings are standardized to 5000 watts when both voltage and current values lie on the power curve.

Table 1. Summary of sampling schedule for rainbow and brown trout mechanical removal trips, 2003-2005.

		FY-	
Trip Type	Trip Date	Year	Trip Length
Electrofishing Depletion	15 – 31 Jan	2003	17 - day
Electrofishing Depletion	12 – 28 Feb	2003	17 - day
Electrofishing Depletion	12 – 28 March	2003	17 - day
Electrofishing Depletion	1- 17 Jul	2003	17 - day
Electrofishing Depletion	1- 17 Aug	2003	17 - day
Electrofishing Depletion	1- 17 Sept	2003	17 - day
Electrofishing Depletion	3 trips Jan-Mar	2004	17 - day
Electrofishing Depletion	3 trips Jul-Sep	2004	17 - day
Electrofishing Depletion	3 trips Jan-Mar	2005	17 - day
Electrofishing Depletion	3 trips Jul-Sep	2005	17 - day
Electrofishing Depletion	3 trips Jul-Sep	2006	17 - day
Electrofishing Depletion	3 trips Jul-Sep	2006	17 - day

Table 2. Description of tasks by day for a typical trip.

	1	non of tasks by day for a typical trip.	Boa	at Electrofis	shing Locat	ions
Trip	Depletion					
Day	Pass	Description of Tasks	Boat 1	Boat 2	Boat 3	Boat 4
1	Control Reach	am: Travel from Lees Ferry to Control Reach Camp. The four electrofishing crews mark sampling unit boundaries. pm: Electrofish a total of 24 sites within the control reach to estimate CPUE for non-native fishes.	7 sites between RM 44-52	between		between
2		am: Travel from Control Reach Camp to Science Beach Camp. The four electrofishing crews deploy hoopnets at standardized locations and begin marking sampling unit boundaries within the Depletion Reach. pm: Set up camp and processing station.				
		am: Complete marking sampling unit boundaries. Organize and ready all gear for depletion operations. Check and pull hoopnet sets at standardized locations. pm: Upstream crews electrofish the upper portions of reaches A and B. Downstream crews electrofish reach C			Reach C,	F1-F6,
3	1	and the upper portions of reaches E and F.	A1-A9	B1-B9	E1-E6	F-14
		am: Crews process fish carcasses from previous day. Deploy hoopnets at standardized locations. pm: Upstream crews electrofish the lower portions of reaches A and B. Downstream crews electrofish the lower				
4	1	portions of reaches E and F.	A10-A19	B10-B19	E7-E13	F7-F13
5	2	am: Crews process fish carcasses from previous day. Check and pull hoopnet sets at standardized locations. pm: Upstream crews electrofish the upper portions of reaches A and B. Downstream crews electrofish reach C and the upper portions of reaches E and F.		B1-B9	Reach C, E1-E6	F1-F6, F-14
		am: Crews process fish carcasses from previous day. Deploy hoopnets at standardized locations. pm: Upstream crews electrofish the lower portions of reaches A and B. Downstream crews electrofish the lower				
6	2	portions of reaches E and F. am: Crews process fish carcasses from previous day. Check and pull hoopnet sets at standardized locations. pm: Upstream crews electrofish the upper portions of		B10-B19	E7-E13	F7-F13
7		reaches A and B. Downstream crews electrofish reach C		D1 D0		F1-F6,
/	3	and the upper portions of reaches E and F. am: Crews process fish carcasses from previous day. pm: Upstream crews electrofish the upper portions of reaches A and B. Downstream crews electrofish Reach C and the	S .	B1-B9	E1-E6	F-14
8	3	upper portions of reaches E and F.		B10-B19	E7-E13	F7-F13
9	4	am: Crews process fish carcasses from previous day. pm: Upstream crews electrofish the lower portions of reaches A and B. Downstream crews electrofish the lower portions of reaches E and F.	l l	B1-B9	Reach C, E1-E6	F1-F6, F-14
10		am: Crews process fish carcasses from previous day. pm: Upstream crews electrofish the upper portions of reaches A and B. Downstream crews electrofish Reach C and the				
10	4	upper portions of reaches E and F. am: Crews process fish carcasses from previous day. pm: Upstream crews electrofish the lower portions of reaches		B10-B19	E7-E13	F7-F13
11	5	A and B. Downstream crews electrofish the lower portions of reaches E and F.	A1-A9	B1-B9	Reach C, E1-E6	F1-F6, F-14

Table 2 (Continued). Description of tasks by day for a typical trip.

Crew Electrofishing Locations

			CICW LICE	uonsiing i	Jocations	
Trip	Depletion					
Day	Pass	Description of Tasks	Crew 1	Crew 2	Crew 3	Crew 4
12	5	am: Crews process fish carcasses from previous day. pm: Upstream crews electrofish the upper portions of reaches A and B. Downstream crews electrofish Reach C and the upper portions of reaches E and F.		B10-B19	E7-E13	F7-F13
13		am: Crews process fish carcasses from previous day. All data and sample boxes inventoried and packed up for runout. Unnecessary equipment packed up for runout. Move to Tanner Beach Camp.				
14		am: Fisheries biologists and technicians hike out Tanner or Bright Angel Trail depending on conditions. Boatman begin runout				
15-17		Run out. Arrive Diamond Creek on Day 17				

Table 3a. Boat and Driver assignments for electrofishing control operations for the January-March sampling trips.

January

	Boat Driver										
Dier	ker	We	iss	Ber	ger	Reeder					
Boat Type	Sample Sites	Boat Type	Sample Sites	Boat Type	Sample Sites	Boat Type	Sample Sites				
Achilles	Z-3	Achilles	Z-17	Aluminu	Z-31	Aluminu	Z-42				
				m		m					
Smith-	Z-4	Smith- Z-19	Coeffelt	Z-32	Coeffelt	Z-43					
Root		Root									
	Z-6		Z-20		Z-35		Z-56				
	Z-7		Z-23		Z-36		Z-57				
	Z-11		Z-27		Z-37		Z-58				
	Z-13				Z-39						
	Z-14				Z-41						

February

	Boat Driver										
Ree	der	Dier	ker	Ber	ger	Weiss					
Boat Type	Sample Sites										
Aluminum	Z-1	Aluminu	Z-15	Achilles	Z-31	Achilles	Z-45				
		m									
Smith-	Z-4	Smith-	- Z-17	Coeffelt	Z-36	Coeffelt	Z-48				
Root		Root									
	Z-6		Z-20		Z-38		Z-51				
	Z-7		Z-25		Z-39		Z-55				
	Z-10		Z-30		Z-40		Z-57				
	Z-11				Z-43						
	Z-13				Z-44						

March

	Boat Driver									
Ree	der	Bei	rger	We	eiss	Dierker				
Boat Type	Sample Sites	Boat Type	Sample Sites	Boat Type	Sample Sites	Boat Type	Sample Sites			
Achilles	Z-5	Achilles	Z-20	Aluminu m	Z-32	Aluminu m	Z-50			
Smith- Root			Coeffelt	Z-36	Coeffelt	Z-51				
	Z-10		Z-26		Z-38	,	Z-52			
	Z-11		Z-27		Z-43		Z-58			
	Z-14		Z-28		Z-46		Z-60			
	Z-16				Z-47					
	Z-17				Z-49					

Table 3b. Boat and Driver assignments for electrofishing removal operations by depletion pass for the January-March sampling trips.

January

	Depletion Pass ^a						
River Reach (Boat Type-Shock Box)	1	2	3	4	5		
A (Rubber-Smith Root)	Dierker	Reeder	Berger	Reeder	Dierker		
B (Alum- Coeffelt)	Weiss	Berger	Reeder	Dierker	Berger		
E & C (Alum-Smith Root)	Reeder	Dierker	Weiss	Berger	Reeder		
F (Rubber-Coeffelt)	Berger	Weiss	Dierker	Weiss	Weiss		

February

	Depletion Pass ^a						
River Reach (Boat Type)	1	2	3	4	5		
A (Alum-Smith Root)	Berger	Reeder	Dierker	Dierker	Dierker		
B (Rubber-Coeffelt)	Dierker	Berger	Weiss	Reeder	Weiss		
E & C (Rubber-Smith Root)	Weiss	Dierker	Berger	Berger	Reeder		
F (Alum-Coeffelt)	Reeder	Weiss	Reeder	Weiss	Berger		

March

			Depletion P	ass ^a	
River Reach (Boat Type)	1	2	3	4	5
A (Rubber-Smith Root)	Dierker	Berger	Weiss	Berger	Weiss
B (Alum-Coeffelt)	Berger	Weiss	Dierker	Weiss	Dierker
E & C (Alum-Smith Root)	Weiss	Dierker	Reeder	Dierker	Berger
F (Rubber-Coeffelt)	Reeder	Reeder	Berger	Reeder	Reeder

 $^{^{\}rm a}$ Depletion pass 1=Day 3 & 4; Depletion pass 2=Day 5 & 6; Depletion pass 3=Day 7 & 8; Depletion pass 4=Day 9 & 10; Depletion pass 5=Day 11 & 12.

Table 4. Target sample sizes and length groups for diet analysis of each species of trout in three Colorado River study areas: Lee's Ferry, above the Little Colorado River, and below the Little Colorado River, Grand Canyon, Arizona.

Length group	Lees Ferry	Above-LCR	Below-LCR
< 150 mm	30	30	30
151 - 200 mm	30	30	30
201 - 300 mm	30	30	30
>301 mm	30	30	30
TOTAL	120	120	120

Table 5. Logistics outline for invertebrate drift sampling to estimate prey availability in the Colorado River above and below the confluence with the Little Colorado River, Grand Canyon, Arizona.

		Station ID										
Above				Above LCR						LCR		
Trip Day ^a	1 2 3 4 5 6					6	7	8	9	10	11	12
3	4	4	4				4	4	4			
4				4	4	4				4	4	4
Total	4	4	4	4	4	4	4	4	4	4	4	4

^a Trip Days 3 and 4 correspond with the first depletion run.

APPENDIX A. ELECTROFISHING DEPLETION DATA SHEET.

Volts: Comments:				AMPS:					Turbidity:(H/L) Total Seconds:				Boat Type Crew:,		
SP	TL	FL	WT	SEX	CON	F CLIP I	LOT # Or BAG ID	PIT RECAP	PIT-TAG#	TAG2 RECAP	TAG2	DISP	COMMENTS		
				12											
		- Va													
											3.				

APPENDIX B. FISH PROCESSING DATA SHEET.

		LECTROFI	SHING E		Page of								
Trip: STAT		D		n nume		_	STORE BOX #						
SPECIES	TL	WT	GAPE VERT	GAPE LAT	F CLIP	PIT RECAP	PIT-TAG#	SEX M/F	SEX	EMPTY (Y/N)	ISOTOPE SAMPLE (Y/N)	ВОТ-#	COMMENTS
N													
		- 10											

APPENDIX C. STANDARD NETTING DATA SHEET.

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)	TL	FL	WT	SEX	Cond	Char	Par Type	Par #		Clip 1		Clip 2	Lot# Or BAG ID	PIT RECAP	PIT-TAG #	DISP	BOT-#		Comments	
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			100																	
٦																				

Appendix D. Standardized Methods For Handling Fish In Grand Canyon Research. Adapted from Ward (2002).

General Guidelines For Handling Fish During Research

Respectful and careful treatment of fish during research is essential to the long-term success of monitoring programs. Traumatized fish can exhibit abnormal physiological, behavioral and ecological responses that defeat study purposes. Rough or improper handling of fish is a source of stress that can lead to disease and death. Delayed mortality as a result of improper handling is often not immediately seen by researchers but can occur hours or days later. This can cause misleading study results and poor public opinion resulting in loss of permits and cancellation of projects. Researchers should be sensitive to public perception and be prepared to explain sampling activities. All field personnel should be familiar with and strictly adhere to research permit guidelines and limitations. Sampling procedures should leave areas as undisturbed as possible and capture techniques should minimize injury to fish. Although specific fish handling procedures vary from one project to another all sampling should incorporate the following general guidelines:

- 1. Be respectful of all fish regardless of size and species
- 2. Minimize the time that fish are out of the water
- 3. Change water frequently when fish must be held for more than a few minutes or if you see fish surfacing for air. Remember that handling stress increases as water temperature increases
- 4. Don't put more than 8-10 fish in your workup bucket at one time. Leave the rest in a net in the river to avoid stressing fish.
- 5. Be aware that watch straps, lapel badges and jewelry can damage fish
- 6. Do not hold fish tightly around the throat and avoid touching the gills
- 7. Rinse all sunscreen or lotions from hands prior to handling fish
- 8. Always wet hands and equipment such as nets and fish boards before use. Dry hands and equipment cause damage to fish skin by removing coatings that protect fish from disease.
- 9. Equipment such as length boards and scales become hot in the sun and can damage fish if not wetted prior to use.

When sampling with hoop nets, shake nets when removing them from the water. Check carefully for small fish that may have become lodged between the net folds. Fish mistakenly left in nets are a large source of researcher caused mortality. Native species accidentally killed should be documented, preserved in ethanol to be deposited as voucher or teaching specimens

Protocols For Processing Fish

Native fish - Measure Total Length (TL), Fork Length (FL), and weight. Examine each fish for external parasites and sexual characteristics. Fish over 100 mm TL should be scanned for the presence of a PIT tag and any fish over 150 mm TL that do not have an existing tag should be tagged.

Nonnative fish - Measure TL of all fish and examine all salmonids for the presence of an adipose clip. Those salmonids that are adipose clipped should be examined for the presence of a pit tag.

Length Measurement

Total Length (TL) – Measure from anterior most part of the fish to the tip of the longest caudal fin ray with the lobes of the caudal fin compressed together

Fork Length (FL) – Length from the most anterior part of the fish to the tip of the median caudal fin ray.

Pit Tagging

Passive integrated transponder (PIT) tags allow long-term unique marking of individual fish. Location of PIT tag insertion varies by species.

PIT Tagging humpback chub

- 1. Verify that needle is sharp and clean (Biomark guidelines recommend that needles be changed every 20 fish)
- 2. Sterilize the needle and tag in Ethanol or Isopropyl alcohol
- 3. Hold the fish with the abdomen up and the tail pointing toward you
- 4. Insert the needle just posterior to the pelvic fin (See figure 5)
- 5. The insertion should be on the abdomen of the fish to the right of the mid-ventral line with the tag placed under the left pelvic girdle. The forward position of the pelvic fins on humpback chub allows the tag to be inserted higher on the abdomen than on other species.
- 6. The depth of penetration of the needle should be deep enough to place the tag within the body cavity and as far away for the needle hole as is feasible to prevent tag loss (preliminary data for trout suggests tag loss may be as high as 10% for tags that are injected too shallow.

(Adapted from Biomark guidelines)

PIT tagging suckers

Use the same procedures as for humpback chub with the following exception. Tags should be inserted toward the tail of the fish under the left pelvic girdle of the fish. The needle should be directed posterior so the tag is injected away from the heart and other vital organs.

Verifying Pit Tag Numbers

The error rate when transcribing and entering PIT tag numbers is very high. The following procedures help to minimize errors that occur when transcribing and entering pit tag numbers.

- 1. Verify that the scanner is in Scan Store mode and says "working" on the display when the trigger is pulled. If scanner is not in Scan Store mode press the menu button several times.
- 2. Scan the fish
- 3. Read and record the entire 10-digit code using words instead of letters to avoid
- 1. Confusion of letters and numbers that sound alike.
- 2. Example 12A3F45E6B Read: one, two, alpha, three, fox, four, five, echo, six, bravo
- 3. Always cross zeros when recording PIT tag numbers. This distinguishes a zero from a "D" in the database. When recording PIT tags draw a horizontal line above any letters in the PIT tag number. This will help us distinguish letters from numbers that can often be confused (B and 8, D and 0, S and 5, etc).
- 4. The data recorder repeats the number back to verify that it has been recorded correctly

Clipping Fins

An adipose clip on brown trout are being used as a secondary mark for newly PIT tagged fish to evaluate tag loss. Marking fish by clipping pelvic fin allows population estimates to be made on fish too small to PIT tag effectively. All researchers must be aware of and look for all possible marks (See fin Clip codes).

- 1. Dorsal punch Use caution to avoid ripping dorsal fin when removing the fin punch.
- 2. Pelvic fin clip Remove a majority of the pelvic fin, but the base of the fin must remain intact so regeneration will occur
- 3. Adipose clip Clip at base removing entire adipose fin.

Fin Clip Codes and Locations

RP1 = right pectoral

RP2 = right pelvic

LP1 = left pectoral

LP2 = left pelvic

ADP = adipose

UCD = upper caudal

LCD = lower caudal

DOR = dorsal

ANL = anal

Guidelines For Recording Data

- 1. If you don't record the data in the field, it is highly unlikely it will be reconstructed in the office. It is better to write down too much information.
- 2. Recording data in the field is one of the most important aspects of the research. You can only work as fast as your data taker can record legible data. If you go too fast processing fish, the data gets sloppy. If we are unable to read your handwriting the data is essentially lost. Keep an eye on your data recorder and ask if you are going too fast. Data recorders, please STOP the fish processor and tell them if they are going to fast.
- 3. Do not forget to write Y or N in the recap field when pit tagging or checking for a pit tag.

Guidelines For Filling Out Data Forms

Three data forms will be used to collect data related to fisheries sampling: 1) Depletion Electrofishing Effort, 2) Depletion Electrofishing Effort Processing, and 3) Netting and Trapping Effort. The Depletion Electrofishing Effort forms will be used to record electrofishing sample and fish catch specific information during both the control and removal sampling. The Depletion Electrofishing Effort Processing Form will be used to record all information about the non-native fish removed during the removal sampling. Finally, the Netting and Trapping Effort form will be used to record the data associated with mainstem hoopnetting activities downstream from the confluence of the LCR.

Depletion Electrofishing Effort Form Instructions (Control Reach)

Trip ID: GC plus Year and Month and date trip **started** (yyyy/mm/dd); e.g.

GC20030115

Start Date: Date the electrofishing sample began (mm/dd/yy)

Start Time: Time the electrofishing sample began; military format (hhmm)

Station ID: Name of sampling station; e.g. Z-07

Depletion Number: Leave blank during control section sampling.

Turbidity: Either High (H) or low (L); see code sheet. Boat Type: Either Achilles (ACH) or Aluminum (ALU)

Volts: Read off CPS unit Amps: Read off CPS unit

Total Seconds: Total amount of time spent electrofishing, read off CPS Unit.

Crew: Boatman and Crew initials (Boatman's Initials first)

Comments: Any noteworthy issues regarding the sample.

SP: Species of fish; see code sheet

TL: Total Length; recorded to nearest mm for all fish captured FL: Fork Length; recorded to nearest mm for **native fish only**

Wt: Weight, recorded to accuracy of scale, measure weight only **for native**

fish larger than 50 mm

Sex: Only record for fish expressing gametes. If you do not try to

determine sex, leave field blank. If you try to determine sex but are

unsuccessful, record as U. See code sheet.

Condition: Sexual Condition. Leave blank if you do not attempt to determine

sexual condition, otherwise use code sheet.

F Clip 1 Finclip 1. First column is recapture status of finclip (Y if fish was

captured with a clip, N otherwise). Only record recapture status if you examine a fish for a clip, otherwise leave blank. Only examine the following species for finclip recapture status: RBT, BNT, HBC; leave this field blank for all other species. Second column is type of fin clip; leave blank if no finclip, otherwise see code sheet. All RBT and BNT fitted with a floy tag will also be given a LEFT

PELVIC CLIP (LP2).

Lot# or Bag ID: Pit tag Lot#; record only for fish receiving a **NEW PIT TAG.** Lot # is

printed on the tape strip or bag containing the pit tags.

PIT Recap: Recapture status of any fish containing a pit tag. It is extremely

important that this field be filled out properly. If a fish contains a PIT Tag upon capture, this field should be "Y". If a fish is found not to contain a PIT TAG or is injected with a new PIT Tag, this field should be "N". Scan all native fish >=120 mm and all BNT with an adipose clip for the presence of a PIT Tag. Inject all untagged native fish >=150 mm with a new PIT TAG. This field should be left blank for all native fish < 120 mm, all BNT without an adipose

finclip, and all other fish.

PIT-TAG #: PIT TAG Number of all fish containing a PIT TAG upon release.

Follow *Verifying PIT TAG Numbers* protocols as described above.

Tag2 Recap: Recap status of other tags besides PIT TAGS. This field will be used

to track the application and recapture of floy tags placed on RBT and BNT in the control reach. This field should be "Y" if a fish is captured with a floy tag. If a fish is found without a floy tag or is fitted with a floy tag, this field should be "N". **This field should be blank for all**

species except RBT and BNT.

Tag2: Floy Tag number. Record either recaptured or new floy tag numbers

in this field. Format is: USGS0001.

Disposition: Disposition of fish after sampling. Except for accidental mortalities,

this field should be "RA" in the control reach. Otherwise, see code

sheet

Comment: Comments specific to a particular fish (e.g. Blind in left eye).

Depletion Electrofishing Effort Form Instructions (Removal Reach)

Trip ID: GC plus Year and Month and date trip **started** (yyyy/mm/dd); e.g.

GC20030115

Start Date: Date the electrofishing sample began (mm/dd/yy)

Start Time: Time the electrofishing sample began; military format (hhmm)

Station ID: Name of sampling station; e.g. A-10 or F-02

Depletion Number: Depletion Run Number (1-5).

Turbidity: Either High (H) or low (L); see code sheet. Boat Type: Either Achilles (ACH) or Aluminum (ALU)

Volts: Read off CPS unit Amps: Read off CPS unit

Total Seconds: Total amount of time spent electrofishing, read off CPS Unit.

Crew: Boatman and Crew initials (Boatman's Initials first)

Comments: Note location where net bag containing non-native fish was left and

any noteworthy issues regarding the sample.

SP: Species of fish; see code sheet. Should only contain native fish as all

non-native fish will be processed at the processing station

TL: Total Length; recorded to nearest mm for all fish captured FL: Fork Length; recorded to nearest mm for **native fish only**

Wt: Weight, recorded to accuracy of scale, measure weight only **for native**

fish larger than 50 mm

Sex: Only record for fish expressing gametes. If you do not try to

determine sex, leave field blank. If you try to determine sex but are

unsuccessful, record as U. See code sheet.

Condition: Sexual Condition. Leave blank if you do not attempt to determine

sexual condition, otherwise use code sheet.

F Clip 1 Finclip 1. First column is recapture status of finclip (Y if fish was

captured with a clip, N otherwise). Only record recapture status if you examine a fish for a clip, otherwise leave blank. Only examine the following HBC for finclip recapture status; leave this field blank for all other species. Second column is type of fin clip; leave blank if

no finclip, otherwise see code sheet.

Lot# or Bag ID: Pit tag Lot#; record only for fish receiving a **NEW PIT TAG.** Lot # is

printed on the tape strip or bag containing the pit tags.

PIT Recap: Recapture status of any fish containing a pit tag. It is extremely

important that this field be filled out properly. If a fish contains a PIT Tag upon capture, this field should be "Y". If a fish is found not to contain a PIT TAG or is injected with a new PIT Tag, this field should be "N". Scan all native fish >=120 mm for the presence of a PIT Tag. Inject all untagged native fish >=150 mm with a new PIT TAG. This field should be left blank for all native fish < 120

mm and all other fish.

PIT-TAG #: PIT TAG Number of all fish containing a PIT TAG upon release.

Follow *Verifying PIT TAG Numbers* protocols as described above.

Tag2 Recap: This field should not be needed during removal operations since

non-native fish will be processed at the processing station. Leave

Blank

Tag2: This field should not be needed during removal operations since

non-native fish will be processed at the processing station. Leave

Blank

Disposition: Disposition of fish after sampling. Except for accidental mortalities,

this field should be "RA" in the removal reach since only native fish will be processed by electrofishing crews. Otherwise, see code

sheet.

Comments specific to a particular fish (e.g. Blind in left eye).

Netting and Trapping Effort Data Forms (Mainstem Hoopnetting)

Gear Type: HS (hoopnet small)

Crew: Boatman and Crew initials (Boatman's Initials first)

Clipboard: Leave Blank Ortho Cov: Leave Blank

Trip: GC plusYear and Month and date trip **started** (yyyy/mm/dd); e.g.

GC20030115

Station ID: Refer to Map of Hoopnetting sites. Format is HS-01 through HS-30

River Guide: Leave Blank
River: Leave Blank
Depletion Number: Leave Blank
Water Temp: Leave Blank

Turbidity: Either High (H) or low (L); see code sheet.

Set #: Same as Station ID.

Haul: A, B, or C. A for first set, B for second set, C for third set.

Start Date: Date the net was set (mm/dd/yy)

Start Time: Time the net was set; military format (hhmm)

Start Date: Date the net was checked (mm/dd/yy)

Start Time: Time the net was checked; military format (hhmm)

River Mile: Leave Blank Side: Leave Blank Waypoint: Leave Blank

ShoreHab: Shoreline Habitat; see code sheet HydrUnit: Hydraulic Unit; see code sheet

Substrate: Leave Blank

Covertype: Cover Type; see code sheet

Set Depth: Leave Blank

SP: Species of fish; see code sheet.

TL: Total Length; recorded to nearest mm for all fish captured FL: Fork Length; recorded to nearest mm for **native fish only**

Wt: Weight, recorded to accuracy of scale, measure weight only **for native**

fish larger than 50 mm

Sex: Only record for fish expressing gametes. If you do not try to

determine sex, leave field blank. If you try to determine sex but are

unsuccessful, record as U. See code sheet.

Condition: Sexual Condition. Leave blank if you do not attempt to determine

sexual condition, otherwise use code sheet.

Char: Sexual Characteristics. Leave blank if you do not attempt to

determine sexual condition, otherwise use code sheet.

Par Type: Parasite Type; Leave blank if you do not examine fish for parasites,

otherwise see code sheet.

Par #: Number of parasites.

F Clip 1 Finclip 1. First column is recapture status of finclip (Y if fish was

captured with a clip, N otherwise). Only record recapture status if you examine a fish for a clip, otherwise leave blank. Only examine HBC for finclip recapture status; leave this field blank for all other species. Second column is type of fin clip; leave blank if no finclip,

otherwise see code sheet.

F Clip 2 Finclip 2. First column is recapture status of finclip (Y if fish was

captured with a clip, N otherwise). Only record recapture status if you examine a fish for a clip, otherwise leave blank. Only examine HBC for finclip recapture status; leave this field blank for all other species. Second column is type of fin clip; leave blank if no finclip,

otherwise see code sheet.

Lot# or Bag ID: Pit tag Lot#; record only for fish receiving a **NEW PIT TAG.** Lot # is

printed on the tape strip or bag containing the pit tags.

PIT Recap: Recapture status of any fish containing a pit tag. It is extremely

important that this field be filled out properly. If a fish contains a PIT Tag upon capture, this field should be "Y". If a fish is found not to contain a PIT TAG or is injected with a new PIT Tag, this field should be "N". Scan all native fish >=120 mm for the presence of a PIT Tag. Inject all untagged native fish >=150 mm with a new PIT TAG. This field should be left blank for all native fish < 120

mm and all other fish.

PIT-TAG #: PIT TAG Number of all fish containing a PIT TAG upon release.

Follow *Verifying PIT TAG Numbers* protocols as described above.

Disposition: Disposition of fish after sampling. See code sheet.

Bot-#: Bottle Number. If a tissue sample (e.g. a whole fish or a portion of a

fish) is taken, record bottle number that contains the tissue.

Comment: Comments specific to a particular fish (e.g. Blind in left eye).

Depletion Electrofishing Effort PROCESSING Form Instructions (Removal Reach)

Trip ID: GC plus Year and Month and date trip **started** (yyyy/mm/dd); e.g.

GC20030115

Start Date: Date the electrofishing sample began (mm/dd/yy), read off Data Info

Card

Station ID: Name of sampling station; e.g. A-10 or F-02, read off Data Info Card

Depletion Number: Depletion Run Number (1-5), read off Data Info Card

Total Seconds: Total amount of time spent electrofishing, read off Data Info Card.

Store Box #: The storage box number which contains stomach samples associated

with this data sheet.

Proc. Comments: Any comments relative to all the samples processed on this sheet.

Species:

TL Total Length; recorded to nearest mm for all fish captured

Wt Weight, recorded to accuracy of scale.

Gape Vert: Gape Vertical; Gape size from lower jaw to upper jaw with mouth

opened fully. Do not distort mouth past normal maximum opening to

take this measurement; record to nearest mm.

Gape Lateral; maximum distance between left and right maxilla. Do

not distort mouth past normal maximum opening to take this

measurement; record to nearest mm.

F Clip 1 Finclip 1. First column is recapture status of finclip (Y if fish was

captured with a clip, N otherwise). Only record recapture status if you examine a fish for a clip, otherwise leave blank. Only examine RBT and BNT for finclip recapture status; leave this field blank for all other species. Second column is type of fin clip; leave blank if no

finclip, otherwise see code sheet.

PIT Recap: Recapture status of any fish containing a pit tag. It is extremely

important that this field be filled out properly. If a fish contains a PIT Tag upon capture, this field should be "Y". If a fish is found not to contain a PIT TAG, this field should be "N". Scan all BNT with an adipose clip for the presence of a PIT Tag. This field should be

left blank for all fish except BNT with an adipose clip.

PIT-TAG #: PIT TAG Number of all fish containing a PIT TAG upon release.

Follow *Verifying PIT TAG Numbers* protocols as described above.

Sex Sex of fish as determined through dissection, see code sheet.

Sexual Condition: Sexual condition of fish as determined through dissection, see code

sheet.

Empty: Y if stomach is empty, N otherwise.

Isotope Sample: Y if a stable isotope is taken, N otherwise

Bottle Number: Number of Metal tag placed in the sample bottle and written on the lid.

Comments: Any comments specific to the fish specimen

APPENDIX. E DATA CODES

Fish species codes

Species Codes	
BBH	Black Bullhead
BGS	Bluegill
BHS	Bluehead Sucker
BKC	Black Crappie
BKT	Brook Trout
BNT	Brown Trout
CCF	Channel Catfish
CRP	Carp
CUT	Cutthroat Trout
FHM	Fathead Minnow
FMS	Flannelmouth Sucker
FRH	Flannelmouth/Razorback Hybrid
GAM	Gambusia
GSF	Green Sunfish
GSH	Golden Shiner
HBC	Humpback Chub
LMB	Largemouth Bass
NOP	Northern Pike
PKF	Plains Killifish
RBS	Razorback Sucker
RBT	Rainbow Trout
RGK	Rio Grande Killifish
RSH	Red Shiner
RTC	Roundtail Chub
SDS	Sand Shiner
SHR	Shiner
SMB	Smallmouth Bass
SPD	Speckled Dace
STB	Striped Bass
SUC	Un-identified Sucker
TFS	Threadfin Shad
UID	Un-determined Fish
UTC	Utah Chub
WAL	Walleye
YBH	Yellow Bullhead

Turbidity codes

Turbidity	
Н	High secchi (< 0.5m)
L	Low secchi (> 0.5m)

Shoreline habitat codes

Shoreline Habitat				
BE	Bedrock			
TD	Travertine Dams			
CB	Cobble Bar			
CL	Cliff			
DF	Debris Fan			
SB	Sand Bar			
TA	Talus			
BL	Boulder			
LE	Ledge			

Hydraulic unit codes

Hydraulic Units					
BA	Backwater				
ED	Eddy (countercurrent)				
RI	Riffle				
RU	Run				
RA	Rapid				
PO	Pool (still)				
RC	Return Channel				
GL	Glide				

Sexual condition codes

Sexual Condition						
N	Not Ripe					
	Mature, Non-Extrudable Developed					
M	Gamates					
R	Ripe, Gametes Exrudable					
S	Spent, Expelled Gametes					
U	Undetermined					

Sexual characteristics

Sexual Characteristics					
C	Color				
T	Tuberculate				
В	Both Colored and Tuberculate				
U	Undetermined				

Cover type codes

Cover Types						
V	Vegetative					
В	Boulder					
L	Ledge or Lateral Cover					
N	None					
U	Undetermined					

Disposition codes

Disposition	
RA	Returned Alive
DR	Dead, Released
DP	Dead, Preserved
	Dead, Stomach
DS	Contents
SK	Dead, Skeletonized

Fin clip codes

Fin Clips	
(Y/N)	Recap, Fin Mark
ADP	Adipose
LP1	Left Pectoral
LP2	Left Pelvic
RP1	Right Pectoral
RP2	Right Pelvic
UCD	Upper Caudal
LCD	Lower Caudal
ANL	Anal

Parasite codes

Parasite Type	
L	Lernea
A	Asian Tapeworm
U	Undetermined
N	None

APPENDIX F. PERSONNEL FOR MECHANICAL REMOVAL OF NON-NATIVE SALMONIDS.

Trip 1 – January 15-31, 2003

HSS

Logistical Support Boatman
 Logistical Support Boatman
 Logistical Support Boatman
 Logistical Support Boatman
 Electrofishing Boatman

GCMRC

8. Processing/Transport Boatman Michael Yard (out on trip day 5)

9. Processing/Transport Boatman Lew Coggins

10. Processors-Field-hands11. Processors -Field-handsMelanie Caron

12. Netters-Field-hands Ted Kennedy (out on trip day 5)

HSS

13. Netters-Field-handsAlley Martinez14. Netters-Field-handsCourtney Giauque15. Netters-Field-handsDanny Martinez16. Netters-Field-handsYael Bernstein17. Processor/Logistical Support TraineeSteve Jones18. Netters-Field-hands??(Volunteer)

Hualapai Nation

19. Netters-Field-hands Aaron Mapatis